4160-01-P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2013-D-0710]

Draft Guidance for Industry on Circumstances that Constitute Delaying, Denying, Limiting, or Refusing a Drug Inspection; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a draft guidance for industry entitled "Circumstances that Constitute Delaying, Denying, Limiting, or Refusing a Drug Inspection." The Food and Drug Administration Safety and Innovation Act (FDASIA) added a new provision to the Food, Drug, and Cosmetic Act (FD&C Act) concerning inspections that would make a drug adulterated. This guidance defines, by way of example, the circumstances that FDA would consider to constitute delaying, denying, or limiting inspection, or refusing to permit entry or inspection for the purposes of making a drug adulterated.

DATES: Although you can comment on any guidance at any time (see 21 CFR 10.115(g)(5)), to ensure that the Agency considers your comment on this draft guidance before it begins work on the final version of the guidance, submit either electronic or written comments on the draft guidance by [INSERT DATE 60 DAYS AFTER DATE OF PUBLICATION IN THE FEDERAL REGISTER].

ADDRESSES: Submit written requests for single copies of the draft guidance to the Office of Policy and Risk Management, Office of Regulatory Affairs, Food and Drug Administration, 12420 Parklawn Dr., rm. 4138, Rockville, MD 20857. Send one self-addressed adhesive label

to assist that office in processing your requests. See the SUPPLEMENTARY INFORMATION section for electronic access to the draft guidance document.

Submit electronic comments on the draft guidance to <a href="http://www.regulations.gov">http://www.regulations.gov</a>.

Submit written comments to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT: Emily M. Leongini, Office of Policy and Risk Management, Office of Regulatory Affairs, Food and Drug Administration, 10903 New

Hampshire Ave., Bldg. 32, rm. 4339, Silver Spring, MD 20903, 301-796-5300.

## SUPPLEMENTARY INFORMATION:

## I. Background

FDA is announcing the availability of a draft guidance for industry entitled "Circumstances that Constitute Delaying, Denying, Limiting, or Refusing a Drug Inspection." On July 9, 2012, FDASIA (Public Law 112-144) was signed into law. Section 707 of FDASIA adds 501(j) to the FD&C Act (21 U.S.C. 351(j) to make a drug adulterated that "has been manufactured, processed, packed, or held in any factory, warehouse, or establishment and the owner, operator, or agent of such factory, warehouse, or establishment delays, denies, or limits an inspection, or refuses to permit entry or inspection." As required by section 707, FDA is issuing this guidance to define the types of action, inaction, and circumstances that FDA considers to constitute delaying, denying, or limiting inspection, or refusing to permit entry or inspection for the purposes of section 501(j) of the FD&C Act.

This draft guidance is being issued consistent with FDA's good guidance practices regulation (21 CFR 10.115). The draft guidance, when finalized, will represent the Agency's current thinking on "Circumstances that Constitute Delaying, Denying, Limiting, or Refusing a

3

Drug Inspection." It does not create or confer any rights for or on any person and does not

operate to bind FDA or the public. An alternative approach may be used if such approach

satisfies the requirements of the applicable statutes and regulations.

II. Comments

Interested persons may submit either electronic comments regarding this document to

http://www.regulations.gov or written comments to the Division of Dockets Management (see

ADDRESSES). It is only necessary to send one set of comments. Identify comments with the

docket number found in brackets in the heading of this document. Received comments may be

seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through

Friday, and will be posted to the docket at <a href="http://www.regulations.gov">http://www.regulations.gov</a>.

III. Electronic Access

Persons with access to the Internet may obtain the document at either

http://www.fda.gov/RegulatoryInformation/Guidances/ucm122044.htm or

http://www.regulations.gov.

Dated: July 9, 2013.

Leslie Kux,

Assistant Commissioner for Policy.

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